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PATENT AND TRADEMARK OFFICE 2292/0H795

PRIORITY DATE CLAIMED

30 April 1998

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DESIGNATED/ELECTED OFFICE (DO/EO/US) INTERNATIONAL APPLICATION NO.

INTERNATIONAL FILING DATE

30 April 1999

PCT/GB99/01350

TRANSMITTAL LETTER TO THE UNITED STATES

TITLE OF INVENTION

IMMUNOSUPPRESSION BY BLOCKING T CELL CO-STIMULATION SIGNAL 2(B7/CD28 INTERACTION)

APPLICANT(S) FOR DO/EO/US

Ian Robert LECHLER and Anthony DORLING

Applicant herewith submits to the United States Designated/Elected office (DO/EO/US) the following items and other information:

- 1. [X] This is a FIRST submission of items concerning a filing under 35 U.S.C. 371.
- This is a SECOND or SUBSEQUENT submission of items concerning a filing under 35 U.S. C. 371. 2. []
- This is an express request to begin national examination procedures (35 U.S.C. 371 (f)) at any time rather than delay examination until 3. [] the expiration of the applicable time limit set in 35 U.S. C. 371 (b) and PCT Articles 22 and 39 (1).
- A proper Demand for International Preliminary Examination was made by the 19th month from the earliest claimed priority date. 4. [X]
- 5. [Xt A copy of the International Application as filed (35 U.S. C. 371 (c) (2))
 - a. [] is transmitted herewith (required only if not transmitted by the International Bureau).
 - b. [X] has been transmitted by the International Bureau
 - c. [] is not required, as the application was filed in the United States Receiving Office (RO/US)
- A translation of the International Application into English (35 U.S. C. 371 (c)2)). 6. [] ₹
- 7. 0 Amendments to the claims of the International Application under PCT Article 19 (35 U.S.C. 371 (c)(3)) a. [] are transmitted herewith (required only if not transmitted by the International Bureau).
 - b. [] have been transmitted by the International Bureau.
 - c. [] have not been made; however, the time limit for making such amendments has NOT expired.
 - d. [] have not been made and will not be made. in
- A translation of the amendments to the claims under PCT Article 19 (35 U.S.C. 371 (c) (3)). 8. E.
- An oath or declaration of the inventor(s) (35 U.S.C. 371(c)(4)) (unsigned).
- A translation of the annexes to the International Preliminary Examination Report under PCT Article 36 (35 U.S.C. 371 (c)(5)). 10. 11

Items 11. to 16. below concern other document(s) or information included:

- An Information Disclosure Statement under 37 CFR 1.97 and 1.98 (with ^C references). 11. []
- An assignment document for recording. A separate cover sheet in compliance with 37 CFR 3.28 and 3.31 is included. 12.[]
- 13. [X] A FIRST preliminary amendment.
 - A SECOND or SUBSEQUENT preliminary amendment. []
- 14. [] A substitute specification.
- A change of power of attorney an/or address letter. 15.[]
- 16. [] Other items or information:

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Ų,S, APPLICATION NO. (If K	nown sec 37 C.F.R.1.50)	INTERNATIONAL APPLICATION			ocket Number
17. [x] The following fees	ALCULATIONS	PTO USE ONLY			
Basic National Fee (37 C Search Report has been	FR 1.492 (a)(1)-(5)): prepared by the EPO [X] or .	PO []	\$860.00		
International preliminary examination fee paid to USPTO (37 CFR 1.482)					
No international preliminary examination fee paid to USPTO(37 CFR 4.482) but international search fee paid to USPTO (37 CFR 1.445 (a) (2) \$710.00					
Neither international pre 1.445(a)(2)) paid to USF		CFR 1.482) nor international se	earch fee (37 CFR \$1,000.00		
International preliminary examination fee paid to USPTO (37 CFR 1.482) and all claims satisfied provisions of PCT Article 33(2)-(4)					
ENTER APPROPRIATE BASIC FEE AMOUNT =					60.00
Surcharge of \$130.00 for furnishing the oath or declaration later than []20 []30 months from the earliest claimed priority date (37 CFR 1.492(e)).					
Claims	Number Filed	Number Extra	Rate		
Total Claims	30-20	10	10 X \$18.00	\$180.00	
Independent Claims	6-3	3	3 X \$80.00	\$240.00	
Multiple dependent claims(s) (if applicable) +270					
TOTAL OF ABOVE CALCULATIONS =					
Reduction by 1/2 for filing by small entity, if applicable. Verified Small Entity statement must also be filed. (Note 37 GFR 1.9, 1.27, 1.28).					
100			SUBTOTAL =	\$1,280.00	
Processing fee of \$130.00 for furnishing the English translation later the [] 20 [] 39 months from the earliest claimed priority date (37 CPR 1.492(f)). +				\$	
to to			TOTAL NATIONAL FEE =	\$1,280.00	
Fee for recording the enclosed assignment (37 CFR 1.21(h)), the assignment must be accompanied by an					

[X] A check in the amount of \$1,280.00 to cover the above fees is enclosed.

appropriate cover sheet (37 CFR 3.28, 3.31). \$40.00 per property

- b. [] Please charge my Deposit Account No.04-0100 in the amount of \$ to cover the above fees.
- c. [X] The Commissioner is hereby authorized to charge any additional fees which may be required, or credit any overpayment to Deposit Account No. 04-0100. A duplicate copy of this sheet is enclosed.

NOTE: Where an appropriate time limit under 37 CFR 1.494 or 1.495 has not been met, a petition to revive (37 CFR 1.137(a) or (b)) must be filed and granted to restore the application to pending status.

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SIGNATURE Faul Fellows

TOTAL FEES ENCLOSED =

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NAME Paul F. Fehiner

REGISTRATION NO. 35,135

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2292/0H795

IN THE UNITED STATES PATENT AND TRADEMARK OFFICE

In re Application of:

Ian Robert LECHLER and Anthony DORLING

Serial No: To Be Assigned

(U.S. National Phase of International Application No. PCT/GB99/01350 - filed 30 April 1999)

Filed: Concurrently herewith

For: IMMUNOSUPPRESSION BY BLOCKING T CELL CO-STIMULATION

SIGNAL 2(B7/CD28 INTERACTION)

Honorable Commissioner of Patents and Trademarks

BOX PCT

Washington, D.C. 20231 ATTN: DO/EO/US

PRELIMINARY AMENDMENT

Sir:

Prior to examination, applicants wish to amend the above-identified application as

follows:



IN THE CLAIMS:

Please amend the claims as follows:

- (Amended) Nucleic acid which encodes a protein according to claim 8 [or claim 9].
- (Amended) A cell which expresses a protein according to claim 8 [or claim 9]
 on its surface.
- (Amended) An animal comprising a cell according to claim 11 [and/or biological tissue according to claim 12].
- 15. (Amended) A process for rendering biological tissue suitable for xenotransplantation, comprising the step of treating said biological tissue such that it expresses a protein according to claim 8 [or claim 9] on the surface of its cells.
- 16. (Amended) A protein according to claim 8 [or claim 9, or of nucleic acid according to claim 10,] in the preparation of a formulation for administration to a xenotransplant recipient or donor.

- 17. (Amended) The use of a protein according to claim 8 [or claim 9, or of nucleic acid according to claim 10,] in the preparation of a formulation for administration to a xenotransplant recipient or donor.
- 20. (Amended) A cell according to claim 18 [or claim 19], wherein said cell does not express B7 on its surface.
- 21. (Amended) A cell according to claim 18 (or claim 19], wherein said cell is a transected immature dendritic cell.
- 22. (Amended) Biological tissue comprising a cell according to [any one of claims 18, 19, 20 or 21] claim 18.
- 26. (Amended) A cell according to [any one of claims 18, 19, 20 or 21,] <u>claim 18</u> for use as a medicament.

Please add claims 29 and 30:

- -- 29. A biological tissue according to claim 12
- 30. A nucleic acid according to claim 10 in the preparation of a formulation for administration to a xenotransplant recipient or donor. --.

REMARKS

The claims have been amended to eliminate multiple claim dependencies.

Claims 29 and 30 have been added.

Entry of this amendment is respectfully requested.

Respectfully submitted,

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Registration No. 35,135

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IMMUNOSUPPRESSION BY BLOCKING T CELL CO-STIMULATION SIGNAL 2 (B7/CD28 INTERACTION)

This invention relates to the suppression of xenograft rejection.

BACKGROUND TO THE INVENTION

The success of allogeneic organ transplantation has been established in the last few decades, but the limited supply of donor organs means that many patients have little or no chance of receiving a transplanted organ, such as a kidney, heart or liver. A significant number of these people die whilst awaiting an organ. One potential solution is "xenografting", or the use of organs from a non-human ("xenogeneic") animal donor.

Porcine donor organs are thought to be suitable candidates because pigs are anatomically and physiologically similar to humans and are in abundant supply. Porcine organs are rejected rapidly upon revascularisation, however, by a humoral process called hyperacute rejection (HAR). This is caused by naturally-occurring antibodies in the recipient which recognise and cross-react with antigens on the endothelial cells (ECs) of the xenograft. This recognition triggers the complement cascade which in turn leads to rejection.

European patent 0495852 (Imutran) suggests that membrane-bound regulators of host complement should be expressed on the xenograft in order to prevent the complete activation of complement in the organ recipient. This approach has been developed and applied in order to produce transgenic animals with organs designed to survive hyperacute rejection [eg. refs 1 & 2].

However, organs surviving HAR are subject to delayed xenograft rejection (DXR). This is characterised by the infiltration of recipient inflammatory cells and thrombosis of graft vessels, leading to ischaemia. WO98/42850 shows that expression of coagulation inhibitors on the surface of the xenograft can inhibit the thrombotic aspect of this type of rejection.

HAR and DXR are followed by the host T lymphocyte-mediated response. There are two pathways, "direct" and "indirect" by which T-cells may become sensitised against xenoantigens. The direct pathway involves interactions between T-cells and MHC molecules on xenogeneic donor cells, whereas the indirect pathway involves the presentation of processed xenoantigens by host APCs in the context of MHC class II. The indirect T-cell response is much stronger against xenoantigens than against alloantigens [3], which contrasts with findings for the direct pathway [4], indicating that both the direct and indirect human T-cell responses against xenoantigens must be suppressed if xenotransplantation is to be effective.

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It appears that the suppression of anti-xenograft indirect T-cell responses will be one of the greatest challenges for xenotransplantation [5,6]. Maintaining the level of immunosuppression required to prevent chronic xenograft rejection due to persistent indirect immunogenicity may be unfeasible using conventional systemic immunosuppressive drugs because of the increased the risks of infection and neoplasia [eg. 7]. Clearly, if xenotransplantation is to be clinically successful, methods to promote graft-specific immunosuppression are needed in order to reduce the requirements for systemic therapy.

T-cell activation requires two separate signals. Delivery of signal 1 alone induces a refractory state ("anergy"), defined as the inability to produce IL-2 after subsequent antigenic exposure. For full activation to occur, the cell must be co-stimulated with signal 2.

In vivo, signal 1 is provided by the interaction of the TCR/CD4 complex with either allogeneic MHC or antigenic peptide complexed with self MHC; signal 2 is supplied by the interaction between B7 molecules (B7.1 and B7.2, also known as CD80 and CD86, respectively) on the antigen-presenting cell (APC) and CD28 on the T-cell

Monoclonal antibodies (mAbs) have played a key role in studying T-cell activation. Signal 1 can be supplied by mAbs directed against the TCR/CD3 complex, and mAbs against CD28 can provide signal 2. Indeed, T-cells can be activated by two suitable mAbs, even in the absence of APC. Activation can also be prevented, rather than provided, using mAbs. Signal 2 can be blocked, for instance, using mAbs which block either B7 or CD28.

Signal 2 can also be blocked by using modified forms of CTLA-4, a high-affinity ligand for B7. CTLA-4 is a natural negative regulator of T-cell activation, and B7 binding to CTLA-4 on an activated T-cell antagonises the co-stimulatory signal provided by the B7/CD28 interaction. Soluble forms of CTLA-4, consisting of the extracellular domains of CTLA-4 linked to the constant domain of an antibody, have been constructed [8,9] to block T-cell activation. These molecules ("CTLA4-Ig" or "CTLA4-Fc") behave in a similar way to anti-B7 antibodies and have been used *in vitro* and *in vivo* to prevent the co-stimulatory functions of B7 and thus promote tolerance [10].

Targeting the B7/CD28 interaction to prevent T cell sensitisation to graft antigens in vivo has been shown to be an effective strategy to enhance graft survival. Using CTLA4-Ig, prolonged survival has been obtained in various allograft models [eg. 11] and in a human-to-murine islet xenograft model [12]. In the xenograft model, CTLA4-Ig administration caused full tolerance against the xenoantigens by rendering direct-reactive T cells anergic.

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It is thus an object of the invention to provide means to promote xenograft-specific immunosuppression. In particular, it as an object of the invention to inhibit T-cell-mediated rejection of xenotransplanted organs by preventing the organ recipient's T-cells from mounting an immune response against the organ. More specifically, it is an object to prevent this immune response by inducing anergy in the recipient's T-cells which recognise the xenotransplanted organ, resulting in xenograft-specific T-cell tolerance.

DESCRIPTION OF THE INVENTION

The invention provides methods and biological reagents for inhibiting T-cell mediated rejection of a xenotransplanted organ by blocking the delivery of co-stimulatory signal 2 in order to prevent the activation of xenoreactive T-cells in the recipient.

This is embodied in three aspects, which are illustrated in Figure 1. It will be appreciated that these three aspects can be used in isolation or in various combinations. Furthermore, conventional immunosuppressive techniques may be used alongside those of the invention.

The following should be read in conjunction with the section entitled "Definitions", which begins on page 8.

The first aspect

In a first aspect, co-stimulation by signal 2 is prevented by administration to the organ recipient of a soluble form of CTLA-4 from the xenogeneic donor organism. If, for instance, a pig organ (donor) were being transplanted into a human (recipient), a soluble form of porcine CTLA-4 (see below) would be administered to the human.

Although CTLA-4 from one organism (eg. pig) is able to bind to B7 from another organism (eg. human), the highest avidity is found for allogeneic B7. Whilst soluble CTLA-4 from the donor organism can thus bind to both recipient B7 (on normal cells) and donor B7 (on xenotransplanted cells), it preferentially binds B7 on the xenograft. This results in xenograft-specific immunosuppression, unlike the administration of CTLA-4 from the recipient organism, which would tend to lead to systemic immunosuppression. By blocking the interaction between B7 on the xenogeneic donor cells and CD28 on recipient T-cells, co-stimulatory signal 2 is not delivered to the T-cell of the recipient. Xenoreactive recipient T-cells are therefore rendered anergic.

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The invention thus provides a method of inducing xenotransplant tolerance in an organ recipient, comprising the administration to said recipient of a soluble form of the CTLA-4 protein from the xenogeneic donor organism.

The soluble form of CTLA-4 preferably comprises a fragment of the CTLA-4 from the donor organism which retains the ability to bind B7. This fragment is preferably the complete extracellular domain of CTLA-4.

Preferably, the soluble protein further comprises the constant domain of an immunoglobulin (eg. the C71 chain of IgG1). Preferably, this is from the recipient organism, in order to prevent an immune response against this portion of the molecule.

In a typical embodiment for pig-to-human transplantation, therefore, the soluble protein could comprise the extracellular domain of porcine CTLA-4 fused to a human $C\gamma 1$ sequence.

Soluble forms of CTLA-4 from other organisms are described in, for instance, references 8 (human CTLA-4/human Ig γ 1 constant region) and 9 (murine CTLA-4/human Ig γ 1).

The invention also provides the use of a soluble form of xenogeneic CTLA-4 in the preparation of a medicament for inducing xenograft tolerance in an organ recipient.

The protein may be administered before, during, or after the xenotransplantation. Pre-xenotransplantation administration is most useful when the recipient is undergoing a pre-transplantation immunisation programme involving exposure to xenogeneic cells.

In the context of a pig being the donor organism, the invention provides a protein comprising the amino acid sequence shown in Figure 2 as SEQ ID:1, which is CTLA-4 cloned from porcine cells. This is the preferred form of CTLA-4 for use in the invention. The extracellular domain of this protein is also shown in Figure 2.

The invention also provides nucleic acid which encodes protein SEQ ID:1 (or fragments thereof). This preferably comprises the nucleotide sequence shown in Figure 3 as SEQ ID:2.

25 In addition, the invention provides a vector comprising the nucleic acid of the invention, and a cell transformed with such a vector.

The second aspect

In a second aspect, co-stimulation by signal 2 is antagonised by expressing a ligand for CTLA-4 on the xenogeneic donor cells. This ligand binds to CTLA-4 on activated T-cells of

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the recipient and antagonises the co-stimulatory signal provided by the interaction between donor B7 and recipient CD28. This renders xenoreactive T-cells anergic.

The invention thus provides a membrane-associated protein which can bind to CTLA-4.

This will typically be a chimeric protein (ie. a protein produced by combining regions of different proteins into a single protein) comprising a CTLA-4-binding region and a membrane-association region. In its simplest form, the protein will thus be a fusion protein

By "membrane-associated protein", it is meant that the protein is attached to a cell membrane such that its extracellular domain can bind to CTLA-4. In order to attach the protein to the cell membrane, the protein might comprise a transmembrane sequence from a membrane protein, for instance, or a GPI anchor. A preferred transmembrane sequence is that of CD4 or CD8. Alternatively the protein might include a sequence which enables it to associate extracellularly with a membrane protein without the protein itself being inserted into the cell membrane.

It may also be desirable for the protein to comprise the cytoplasmic domain which is usually associated with said transmembrane regions (eg. the CD8 cytoplasmic domain), such that the protein is targeted to the cell membrane. Similarly, it may be desirable for the protein to comprise the extracellular sequences immediately juxtaposed with the cell membrane (eg. CD4 domains 3 and 4) in order to separate the CTLA-4-binding domain from the cell membrane. Synthetic linkers, such as glycine linkers, can be used for the same purpose.

The CTLA-4-binding domain of the protein preferably comprises an antibody with specificity for CTLA-4. This is preferably a single chain antibody (sFv). It is preferably specific for the CTLA-4 of a recipient organism.

In a typical embodiment, therefore, a protein of the second aspect can comprise a single chain antibody fused via a linker to the transmembrane and cytoplasmic domains of CD8.

The invention also provides nucleic acid which encodes a protein of the second aspect.

25 In addition, the invention provides a vector comprising said nucleic acid of the invention, and a cell transformed with said vector.

The invention also provides a delivery system comprising nucleic acid, and/or vector according to the second aspect of the invention, and means to deliver this material to a target cell.

Furthermore, the invention provides a cell which expresses a protein of the second aspect on its surface, preferably such that the protein can bind to available CTLA-4.

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So that the cell can engage recipient T-cells, the cell preferably also expresses MHC (class I or class II) on its surface. Suitably, therefore, the cell of the invention is a donor professional APC. Because of the antagonistic signal provided by the anti-CTLA-4 protein, however, these professional APC behave functionally as B7-negative cells.

5 The invention also provides biological tissue comprising such a cell.

The invention further provides an animal comprising a cell and/or biological tissue according to the second aspect.

The invention also provides a process for rendering biological tissue suitable for xenotransplantation, comprising the step of treating said biological tissue such that it expresses one or more proteins according to the second aspect on the surface of its cells.

The invention also provides a method of transplantation comprising the step of transplanting biological tissue according to the invention from a donor animal (eg. a pig) into a xenogeneic recipient animal (eg. a human).

In addition, the cells of the invention are suitable for pre-transplantation administration. This results in tolerance being induced in recipient T-cells before the xenograft itself is transplanted. Whilst the cells used in such pre-transplantation regimes should preferably express MHC class II, it will be appreciated that the cells need not be professional APCs.

Furthermore, the invention provides protein or nucleic acid according to the second aspect for use as a medicament.

20 The invention also provides the use of protein, nucleic acid, a vector, or a delivery system according to the second aspect in the manufacture of a formulation for administration to a xenotransplant recipient or donor.

The third aspect

In a third aspect, co-stimulation by signal 2 is prevented by expressing recipient organism MHC class II on the surface of the cells of the xenogeneic donor organ. If, for instance, a pig organ (donor) were being transplanted into a human (recipient), the pig organ would express human MHC class II

Even if direct activation of recipient T-cells is avoided, for instance by utilising one or both of the first two aspects of the invention described above, indirect activation can still occur, involving the processing and presentation of xenoantigens on MHC class II by recipient APC.

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By expressing recipient MHC class II on the cells of the xenogeneic donor, the donor cells will present xenoantigens to a recipient T-cell in the context of self MHC class II. If the donor cells do not express B7, or if B7 is blocked, the xenoreactive recipient T-cell will not receive co-stimulatory signal 2 and will become anergic before the recipient's APCs have an opportunity to present the xenoantigens themselves.

The invention thus provides a cell which expresses on its surface MHC class II of a different organism. Preferably, this is a porcine cell expressing human MHC class II on its surface.

The MHC class II is preferably of the HLA-DR family.

The MHC class II is preferably constitutively expressed on the surface of the cells.

In order to prevent an allogeneic anti-MHC class II response, the MHC class II is preferably tissue-typed for maximum compatibility with the particular recipient. This will typically involve, for instance, ensuring that the HLA-DR expressed on the xenogeneic donor cell should match the HLA-DR of the particular recipient.

To ensure that xenoantigen display within the groove of the MHC class II molecule mirrors that found on professional APC, it is preferred that the cell should also express one or more of the following three proteins, each of which has an important role in antigen processing: invariant chain, HLA-DMα and HLA-DMβ.

The cell preferably does not express co-stimulatory molecules (eg. B7) on its surface. Typically, therefore, the donor cell is not a professional APC. It may, however, be a transfected non-immunogenic APC, such as an immature dendritic cell, which may be B7⁺.

The invention also provides biological tissue comprising a cell according to the third aspect.

The invention further provides an animal comprising a cell and/or biological tissue according to the third aspect.

The invention also provides a process for rendering biological tissue suitable for xenotransplantation, comprising the step of treating said biological tissue such that it expresses xenogeneic MHC class II on the surface of its cells.

Preferably, this process comprises the steps of isolating non-immunogenic cells (ie. cells which cannot provide a co-stimulatory signal, such as B7-negative cells) from a xenogeneic organism and transfecting these cells with HLA-DR. The HLA-DR is preferably tissue-typed for a specific recipient. Furthermore, the cells may also be transfected with other proteins necessary for efficient antigen processing. Examples of suitable non-immunogenic cells include renal tubular

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epithelial cells, which are B7-negative and have been shown to induce tolerance in rodent models of allogeneic transplantation.

The invention also provides a method of transplantation comprising the step of transplanting biological tissue according to the third aspect from a donor animal (eg. a pig) into a xenogeneic recipient animal (eg. a human).

In addition, the cells of the invention are suitable for pre-transplantation administration. This results in tolerance being induced in recipient T-cells before the xenograft itself is transplanted.

Furthermore, the invention provides a cell according to the third aspect for use as a medicament.

The invention also provides the use of a cell or of biological tissue according to the third aspect in the manufacture of a formulation for administering to a xenotransplant recipient.

The invention also provides the use of xenogeneic MHC class II, or nucleic acid encoding xenogeneic MHC class II, in the preparation of a formulation for administering to a xenotransplant donor.

Definitions

As used above, the term "nucleic acid" includes both DNA and RNA, although modified and synthetic nucleic acids are also included, For instance, the nucleic acid may be synthetic (eg. PNA), or may have modified inter-nucleotide linkages (eg. phosphorothioates). Furthermore, the term includes both sense and antisense nucleic acid sequences, as well as double-stranded sequences.

Preferably the nucleic acid comprises sequences suitable for the regulation of expression of protein according to the invention. This expression can preferably be controlled, such as cell-specific control, inducible control, or temporal control.

As used above, the term "vector" signifies a molecule which is capable of transferring nucleic acid to a host cell, and numerous suitable vectors are known in the art.

Preferably the vector is suitable for the production of a transgenic animal. Vectors suitable for the generation of transgenic pigs, for example, are described in references 13, 14, 15, 16 & 17.

As used above, the term "delivery system" refers to means for delivering genetic material to a target cell.

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Certain vectors as described above may also function as suitable delivery systems. Likewise, certain delivery systems may also inherently be vectors, but this is not always the case. For instance, a viral vector can also function as a delivery system, whereas a liposomal delivery system is not a vector. The delivery system may be viral or non-viral. Non-viral systems, such as liposomes, avoid some of the difficulties associated with virus-based systems, such as the expense of scaled production, poor persistence of expression, and concerns about safety. Preferably the delivery system is suitable for use in gene therapy. Numerous appropriate delivery systems are known in the art.

Preferably, the delivery system will be targeted so that molecules according to the invention are taken up by cells suitable for xenotransplantation, or cells which have been transplanted. More preferably the delivery system will be specific for these cells. For example, the delivery system may be targeted to a specific organ, such as the heart or the kidney, or to a specific cell type, such as endothelial cells or professional APC.

To achieve this the delivery system may, for example, be a receptor-mediated delivery system, being targeted to receptors found on target cells. For example, the delivery system may be targeted to receptors found on heart cells, preferably to receptors found exclusively on heart cells, or it may be targeted to receptors found on endothelial cells, preferably to receptors found exclusively on endothelial cells.

The delivery system is preferably suitable for the generation of a transgenic animal. For example, the delivery system may be targeted to a gamete, a zygote, or an embryonic stem cell.

The vectors and delivery systems of the invention can be used to transfect cells to produce cells according to the invention. The transfection can occur in vivo or ex vivo.

The term "biological tissue" as used above includes collections of cells, tissues, and organs. Accordingly the definition includes, for example, fibroblasts, a comea, nervous tissue, a heart, a liver, or a kidney.

Where the second and third aspects of the invention provide "an animal", said animal is preferably suitable for the production of organs for xenotransplantation and/or cells of the invention (eg. cells for pre-xenotransplant administration to xenograft recipients). Preferably the animal is a mammal, and more preferably it is a transgenic pig or a transgenic sheep.

30 The animal might be treated whilst alive such that it comprises transgenic biological tissue (ie. treated by gene therapy). Preferably, a live animal is transfected with a vector according to the

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invention in order to produce a transgenic animal. For example, a vector according to the invention could be specifically delivered to the heart of a pig to produce biological tissue suitable for xenotransplantation.

Alternatively, the animal might be born as a transgenic animal. Many suitable approaches for generating such transgenic animals are known in the art [eg. refs. 18, 19, 20]. For example, direct manipulation of the zygote or early embryo, by microinjection of DNA for instance, is well known, as is the *in vitro* manipulation of pluripotent cells such as embryonic stem cells. Retroviral infection of early embryos has proved successful in a range of species, and adenoviral infection of zona-free eggs has been reported. Transgenesis and cloning of sheep by nuclear transfer has also been described (eg. WO97/07668).

Where the invention provides a process for rendering biological tissue suitable for xenotransplantation, said biological tissue may be so rendered either in vivo or ex vivo. For example, an animal organ may be in vivo transfected with a vector according to the invention, or an organ could be transfected ex vivo before transplantation or in vivo after transplantation.

BRIEF DESCRIPTION OF THE DRAWINGS

The invention will now be described with reference to the attached drawings, in which:

Figure 1 illustrates the three aspects of the invention. "X" represents a xenogeneic cell (or, in the indirect activation pathway, a xenoantigen-presenting recipient APC), and "T" represents a recipient T-cell. In embodiment I, the delivery of co-stimulatory signal 2 is prevented by using a soluble form of CTLA-4. In embodiment II, anti-CTLA-4 is used to antagonise signal 2. In embodiment III, X expresses recipient MHC-II, but does not express B7.

Figure 2 shows the amino acid sequence of pCTLA-4 (SEQ ID NO:1). The following junctions are illustrated by a "*": signal peptide/extracellular domain; extracellular domain/transmembrane domain; transmembrane domain/cytoplasmic domain. An alignment with the human and bovine sequences is also shown. Homologies with pCTLA4 are:

Domain	Human	Bovine	
Signal peptide	67.6%	86.5%	
Extracellular domain	83.8%	84.6%	
Transmembrane domain	96.1%	100%	
Cytoplasmic domain	100%	100%	
Overall	85.2%	89.2%	

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Figure 3 shows a similar alignment, but at the nucleotide level. Homologies are as follows:

Domain	Human	Bovine	
Signal peptide	76%	81.3%	
Extracellular domain	85.2%	86.3%	
Transmembrane domain	92.3%	96.2%	
Cytoplasmic domain	96.5%	97.7%	
Overall	86.1%	88.3%	

Figure 4 shows the amino acid sequence of the pCTLA4-Ig construct. The underlined sequence shows the flexible linker GGSGGAA, which also denotes the junction between pCTLA4 and the IgG1 domains.

- 5 Figure 5 shows the results of flow cytometric analysis of hCTLA4-Ig (ο & □) and pCTLA4-Ig (ο & Δ) binding to human fibroblasts transfected with either human B7 (lower two lines) or porcine B7 (upper two lines).
 - Figure 6 shows the selective inhibition of proliferation by pCTLA4-Ig $(\circ \& \Delta)$ compared to hCTLA4-Ig $(\Box \& \Diamond)$ when co-stimulated by human B7 $(\Box \& \circ)$ or poreine B7 $(\Diamond \& \Delta)$.
- Figure 7 shows the inhibition of human CD4* T cell proliferation by hCTLA4-Ig (n) or pCTLA4-Ig (0) when human (7A) or porcine (7B) cells expressing MHC-class II were used as stimulators in a five day mixed leukocyte reaction.
- Figure 8 shows the nucleotide sequence of an anti-human CTLA-4 sFv. The inferred protein sequence is shown in Figure 9. Figure 10 shows the nucleotide sequences of four anti-murine
 CTLA-4 sFv. The inferred protein sequences are shown in Figure 11. The heavy and light
 - Figure 12 shows the construct encoding the soluble Ig-fusion of the CTLA-4-specific sFv.
 - Figure 13 shows the inhibition of T cell proliferation by cells expressing either an anti-hCTLA-4 sFv (\square) or a control sFv (\bigcirc) .
- 20 Figure 14 shows construct encoding the membrane-bound form of the anti-CTLA-4 sFv.

chains are linked by a serine-glycine linker as indicated in Figures 9 and 11..

Figure 15 shows (A) the nucleotide sequence and (B) the amino acid sequence of human CTLA-4. The start codon is underlined. At position -21, the sequence differs from GenBank sequence L15006, and at position 110 the sequence differs from both L15006 and M74363.

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Figure 16 shows the sequence of cloned human CD8 α . This differs from the GenBank sequence at positions 231 (T \rightarrow G), 244 (A \rightarrow G), 266 (T \rightarrow C), and 437 (T \rightarrow C).

Figure 17 shows the binding of human and murine CTLA4-Ig to IPEC, in order to define clones as B7-negative or B7-positive.

5 Figure 18 shows the binding to transfected cells of HLA-DR-specific mAb L243.

Figure 19 shows the proliferation by human T-cells to HLA-DR-1 transfected IPEC.

Figure 20 shows the results of a human T-cell proliferation assay following 2 days of incubation HLA-DR-1 transfected cells. The X-axis indicates the stimulator cells used in the second step of the proliferation assay. The black bars show results with CD4 T-cells which were primed with B7-positive transfectants; the white bars (hardly visible) show results after priming with B7-negative transfectants. The first graph shows results with cells harvested on day 3; the second graph shows results from a harvest on the sixth day.

Figure 21 shows the proliferation of an APC-dependent, HLA-DR-1 restricted T-cell line raised against IPEC. The stimulator population is indicated on the X-axis.

15 DESCRIPTION OF EMBODIMENTS

Soluble porcine CTLA-4

Porcine CTLA-4 ("pCTLA4") was cloned from PHA-activated pig T cells. RNA was prepared using standard techniques and pCTLA4 was amplified by PCR using primers:

5'-TTGAAGCTTAGCCATGGCTTGCTCTGGA-3' (5' primer)

5'-TAATGAATTCTCAATTGATGGGAATAAAATAAG-3' (3' primer)

The resulting 700bp fragment was sub-cloned into EcoRI/HindIII digested pBluescript, and the nucleotide sequence was determined using the standard T3 and T7 primers. The sequence of a single clone is shown in figure 3, which also shows a comparison with the human and bovine CTLA-4 sequences.

25 The predicted amino acid sequence of pCTLA4 is shown in figure 2, with a comparison with that of human and cattle. Of significance is the predicted amino acid difference at residue 97, which is important in B7 binding, being part of the conserved hexapeptide motif MYPPPY. In pCTLA4, residue 97 is leucine (giving LYPPPY), whereas other species have methionine (although leucine has also been found in bovine CD28 [21]). This important amino acid

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difference is believed to be of key importance to the advantageous differential binding of pCTLA4 to human and pig B7.

The extracellular domain of pCTLA4 was amplified using the 5' primer described above and 5'-CGGTTCTGCAGCACCACCGGAGCCACCATCAGAATCTGGGCATGGTTCTGGATCAATGAC-3'

This amplified from position 484, introduced an 18 base-pair segment encoding a linker 5 GGSGGAA sequence (underlined), and introduced a PstI site (bold) to allow in-frame ligation to the hinge region of human IgG1. The resulting 500bp fragment was sub-cloned into HindIII/PstI digested pBluescript-IgG1 containing genomic DNA encoding intronic sequences and the hinge, CH2, CH3 and 3' untranslated exons of human IgG1 between PstI/NotI sites. The amino acid sequence of the resulting soluble pCTLA4-Ig is shown in figure 4.

Expression of pCTLA4-Ig

The chimeric pCTLA4-Ig DNA sequence was released from pBluescript as a HindIII/BstXI fragment and was sub-cloned into the expression vector pHOOK-3TM (Invitrogen). This was used to transfect DAP.3 or CHO-K1 cells using standard calcium phosphate precipitation . G418-resistant cells were separated using the CaptureTecTM system. These transfected cells were grown in tissue culture flasks until confluent, at which point the medium was changed, and the cells were kept in culture for a further 3 days. At this stage the medium was harvested and perfused through a protein G column. pCTLA-4-Ig was eluted under acid conditions. The concentration of the eluted protein was calculated using ELISA with an antibody directed against human IgG1 and a standard human IgG1 myeloma protein.

The binding characteristics of pCTLA4-Ig were compared to those of human CTLA4-Ig using flow cytometric analysis with human fibroblasts transfected with either human B7-1 or porcine B7-2. For these experiments, the concentration of pig and human CTLA4-Ig were equivalent as assessed by ELISA. As illustrated in figure 5, human and porcine CTLA4-Ig appeared to have similar binding characteristics on human cells expressing porcine B7. Unlike human CTLA4-Ig, however, pCTLA4-Ig failed to bind human B7, implying that pCTLA4-Ig binds preferentially to porcine B7 and is useful as a species-specific reagent.

pCTLA4-Ig was used to inhibit human T cell proliferative responses to a variety of stimulators. In these assays, co-stimulation of the T cell response was provided by either porcine or human B7, expressed either by transfection or naturally on professional APCs. These experiments are demonstrated in figures 6 and 7.

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In the experiments using transfected fibroblast stimulators (expressing HLA class II and either human or pig B7), hCTLA4-Ig inhibited all proliferative responses (Figure 6, \square & 0). In contrast, pCTLA4-Ig only fully inhibited the response when stimulators expressed porcine B7 (\triangle); the proliferative response to cells expressing human B7 was only minimally affected (\bigcirc).

5 In similar experiments, pCTLA4-Ig failed to have a significant inhibitory effect on the proliferative responses to human cells expressing MHC class II and human B7 but did inhibit the response to porcine stimulators (figure 7).

These results highlight the effective inhibitory properties of pCTLA4-Ig when T cell co-stimulatory signals are provided by porcine B7. The failure to prevent T cell proliferation when co-stimulation is mediated by human B7 also demonstrates species-specific action. It can be concluded that pCTAL4-Ig shows species-specific binding to and inhibition of the functional effects of porcine B7, but not human B7.

Properties of pCTLA-4-Ig

The binding characteristics of pCTLA4-Ig to both human and poreine B7-family molecules may be compared to those of hCTLA4-Ig, for example using the following tests:

- (i) flow cytometric analysis of binding to porcine and human APC, and to transfectants expressing porcine or human B7 (see above)
- (ii) quantitative characterisation of binding using BiacoreTM.
- (iii) functional analysis of the effects of CTLA4-Ig on human anti-pig and human allogeneic mixed lymphocyte cultures.
 - (iv) functional assessment of the ability of pCTLA4-Ig to prolong porcine islet xenograft survival after transplantation into B6 mice.

A membrane-associated protein which binds to CTLA-4

A phage display library containing 10¹² semi-synthetic variable sequences was screened using
25 human or murine CTLA4-Ig and a control human IgG1 myeloma protein. The sFv from a
phage displaying differential binding to the human CTLA4-Ig protein after 4 rounds of
screening were isolated and purified using standard techniques. The nucleotide and inferred
amino acid sequences are shown in figures 8, 9, 10, and 11.

The sFv were amplified by PCR using specific primers based on the nucleotide sequences. The 30 distal portions of the primers were based on sequence within pHOOK1. The 5' primer

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contained an Apal site and the 3' primer contained a Sall site, both of which were predicted to be unique. The resulting sFv were sub-cloned into pBluescript for sequencing to determine faithful amplification. The Apal/SalI fragments were then sub-cloned into pHOOK1, where it is flanked upstream by an in-frame signal sequence from the murine Ig K-chain and a haemaggluttinin A epitope sequence, and downstream by two in-frame myc sequences and a transmembrane sequence from the PDGF receptor.

The myc sequences from pHOOK1 were amplified by PCR using the 5' primer 5'-GAGCTGAAACGGGCGGCAGAAC-3', which contains a NotI site (underlined) and the 3' primer 5'-CTGGCCTGCAGCATTCAGATCC-3', which introduced a PstI site (underlined). The resulting 113 base pair fragment was sub-cloned into NotI/PstI digested pBluescript.

The sFv was released from pHOOK1 as an EcoRI/NotI fragment, and was ligated into EcoRI/PstI digested pBluescript-IgG1, along with the NotI/PstI PCR product [Figure 12]. This construct encodes a soluble Ig-fusion of the CTLA-4-specific sFv. For expression in eukaryotic cells, the construct was sub-cloned into pHOOK3 as a HindIII/BstXI fragment.

To confirm cell-surface expression of the sFv, the pHOOK construct was transfected into cells already expressing HLA-DR molecules and human B7. Cells resistant to G418 or mycophenolic acid, depending on the vector used, were grown in culture. Cells expressing the anti-CTLA4-sFv construct on the cell surface were identified by flow cytometric analysis using hCTLA4-Ig. These cells were cloned by limiting dilution and were used as stimulators of T cell proliferation in 5 day cultures. The results of one experiment are shown in figure 13. Cells expressing the anti-hCTLA4 sFv failed to stimulate T cell proliferation (11), whereas those expressing a control sFv stimulated proliferation in the same way as normal cells (0).

In different experiments, the EcoRI/SalI fragment of the construct shown in Figure 12 was co-ligated with the transmembrane and cytoplasmic domains of human CD8 (isolated as a Sall/BamHI fragment from pBluescript-hCD8) into EcoRI/BamHI digested pBluescript [Figure 14].

The EcoRI/BamHI fragment from pBluescript was sub-cloned into the expression vector pHBApr-1-neo or the sister vector pHBApr-1-gpt. These were transfected into cells already expressing HLA-DR molecules and B7 and selected as described above for the pHOOK construct.

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Membrane-associated CTLA-4 construct

The expression of CTLA-4 on by activated T-cells is only transient so, to test the functional characteristics of the anti-CTLA4-sFv, chimeric constructs comprising the DNA sequences encoding the extracellular domains of human or murine CTLA4 and the transmembrane/cytoplasmic sequences of human CD8 were made. Cells expressing these constructs can be used for the study of the anti-CTLA4-sFv protein.

RNA from PHA-activated human T cells was prepared using standard techniques. hCTLA4 was amplified PCR using primers:

- 5'-TTCAAAGCTTCAGGATCCTGAAAGGTTTTG-3' introducing a HindIII site (5' primer)
- 5'-TAATGAATTCTCAATTGATGGGAATAAAATAAG-3' introducing an EcoRI site (3' primer)

The resulting fragment was sub cloned into HindIII/EcoRI digested pBluescript and the nucleotide sequence determined using standard T3 and T7 primers. The sequence of a single clone is shown in figure 15. This differed by a single base (position 439) from GenBank-listed sequences for human CTLA-4. The predicted amino acid sequence of hCTLA4 is also shown.

- 15 The extracellular domain of hCTLA-4 was amplified using 5' primer described above and:
 - $\texttt{5'-GATGTAGATATCACAGGCGAA} \\ \textbf{CCACCGGAGCCACCAATTACATAAATCTGGGCTCCGTTGCCTATGCCC-3'} \\ \textbf{CCC-3'} \\ \textbf$

This amplified from position 457 and included a 15 base segment encoding a flexible GGGGG amino acid linker (underlined), along with a unique SaII site (highlighted). The resulting fragment was sub-cloned into HindIII/SaII digested pBluescript and sequenced.

- 20 hCD8 was PCR-amplified from resting T-cells using primers:
 - 5'-TCGCGCCCAAGCTTCGAGCCAAGCAGCGT-3' introducing a HindIII site (5' primer)
 - 5'-TAATGAATTCTCAATTGATGGGAATAAAATAAG-3' introducing an EcoRI site (3' primer)

The resulting fragment was sub cloned into HindIII/EcoRI digested pBluescript and the nucleotide sequence determined using standard T3 and T7 primers. The sequence of a single clone is shown in figure 16. This clone differed from the sequence deposited with GenBank at four positions, although none of these were within the region that was subsequently amplified.

The transmembrane (TM) and cytoplasmic (C) domains of hCD8 were amplified using the 3' primer described above and the following 5' primer:

 $[\]texttt{5'-CATAGGCAACGGAGCCCAGATTTATGTAATT} \underline{\texttt{GGTGGCTCCGGTGGT}} \mathbf{GTCGAC} \mathbf{TCGCCTGTGATATCTACATC-3'}$

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This amplified from position 532 and included a 15 base segment encoding a flexible GGSGG amino acid linker (underlined), along with a unique SaII site (highlighted). The resulting fragment was sub-cloned into HindIII/SaII digested pBluescript and called pBluescript-hCD8.

The extracellular domain of human CTLA-4 was cut from pBluescript as an *EcoRI/SaII* fragment, and the TM-IC domain of CD8 cut as a *SaII/BamHI* fragment. Together they were ligated back into *EcoRI/BamHI* digested pBluescript. The whole CTLA-4-CD8 chimera was then removed as a single *EcoRI* fragment and was sub-cloned into a number of expression vectors for expression into the human T cell leukaemia line 16.

Properties of the cell-surface anti-CTLA4 proteins

The cell-surface anti-CTLA-4 proteins may be further characterised by the following functional tests:

- Flow cytometric assessment of the interaction between cells expressing the membrane-bound anti-CTLA4-sFv-CD8 protein and soluble human CTLA4-Ig.
- Quantitative assessment of the interaction between the soluble anti-CTLA4sFv-Ig fusion protein and soluble human CTLA4-Ig, using BiacoreTM
- iii) Analysis on the signalling events resulting from the binding of soluble human CTLA4-Ig to J6 transfectants expressing the anti-CTLA4-sFv-CD8 fusion protein.
- iv) Analysis of T cell responses (eg. proliferation, cytokine production, anergy induction) when stimulation in an allogeneic mixed lymphocyte response is provided by an HLA-DR-positive, B7-positive, anti-CTLA4-sFv-CD8-positive cell line.

B7-negative porcine cells expressing murine MHC class II

Fifty cloned immortalised porcine aortic endothelial cells (PAEC) were generated from monolayers of PAEC by intranuclear microinjection with pZipSVU19 DNA [22]. From the immortalised cells (IPEC), B7-negative clones were identified by flow cytometric screening with hCTLA4-Ig and mCTLA4-Ig [see figure 17]. These were then transfected with cDNAs encoding HLA-DRA and DRB1*0101 in the plasmid expression vectors pcExV1-gpt and pHβApr-lneo, and cells were placed under selection with MXH and G418. For comparison, B7-positive IPEC controls were generated similarly [4].

Another series of IPEC transfectants expressing the murine MHC class II molecule I-A^b were also generated for experiments in mice.

Surface expression of MHC class II on transfected IPEC cells was detected using monoclonal antibody L243 (specific for HLA-DR) [figure 18] or M5-114 (specific for murine MHC class II). MHC class II-positive cells underwent several rounds of fluorescence activated cell sorting before being cloned by limiting dilution.

A second batch of transfectants was prepared in exactly the same way, but with additional transfected cDNAs encoding HLA-DMA and HLA-DMB and p31li (invariant chain) in the expression vector pCMU.

Anergy induction in allogeneic T-cells by MHC class II-expressing cells

Human T-cells were purified using standard protocols [3]. For primary proliferation assays, T-cells were incubated for 5 days with fixed numbers of irradiated stimulator cells, before addition of 1µCi ³H-thymidine sixteen hours prior to harvesting onto glass fibre filters. The filters were read in a scintillation counter.

B7-positive IPEC caused significant, anti-DR1 specific proliferative responses, whereas B7-negative IPEC failed to initiate any proliferative response [figure 19].

Two step anergy induction assays were established by a standard protocol [23]. In the primary, tolerance-induction step, T-cells incubated with B7-positive IPEC mounted an anti-DR1 proliferative response in the secondary step with the kinetics of a primed secondary immune response (maximal at three days). However, T-cells incubated with B7-negative IPEC in the primary step became tolerant to DR1 and failed to mount a response on subsequent exposure to DR1-expressed on B7-positive IPEC [figure 20].

Anergy induction in DR1-restricted T cells by DR1-expressing pig cells.

CD4* T-ceils from a DR1-expressing individual were purified according to standard procedures. In primary proliferation assays, they proliferated to B7-positive IPEC transfected with HLA-DR1, indicating that the DR1 can perform as a restriction element for pig-peptide-specific T-ceils. Assays comparing the proliferative response to B7-positive and B7-negative DR1+ transfectants are being performed.

Two step anergy induction assays may also be performed to demonstrate that DR1-transfected, B7-negative pig cells induce anergy in HLA-DR-restricted human T-cells.

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Overlap between the pig peptides processed by professional human APC for presentation on HLA-DR and those presented on MHC class II of IPEC transfected with HLA-DR.

A human T-cell line against wild type IPEC was raised from human PBMC. The proliferative response of this line was dependent on the presence of human APC and inhibitable by antibodies against HLA-DR, indicating that the line had indirect specificity for processed porcine xenoantigens presented by human APC.

This line proliferated against B7-positive HLA-DR1-transfected IPEC [figure 21] implying that at least some of the processed pig peptides presented indirectly by professional human APC are also presented by transfected pig cells.

10 Studies in pig-islets-to-mouse model

In vivo, porcine pancreatic islet cells may be transplanted under the kidney capsule of streptozotocin-treated diabetic mice. Islet xenografts, being non vascular, are rejected solely by T-cells. Porcine islets are prepared from the pancreas of pigs under terminal anaesthesia, and their survival in the recipients assessed by maintenance of normoglycaemia. Mice are injected intravenously with B7-negative, I-A^b-expressing IPEC before transplantation of pig islets. This strategy can be combined with other aspects of the invention to tolerise the direct pathway of T-cell recognition, to ensure that rejection via the direct pathway does not occur. To assess whether a particular strategy has induced specific T-cell tolerance, nephrectomy of the islet-carrying kidney is performed before re-transplantation (under the capsule of the surviving kidney), of identical or third party porcine islets.

It will be understood that the invention is described above by way of example only and modifications may be made whilst remaining within the scope and spirit of the invention.

REFERENCES (The contents of which are incorporated in full)

- 1 Squinto SP (1996) Xenogeneic organ transplantation. Curr Opin Biotech 7: 641-645.
- 2 McCurry et al. (1996) Human complement regulatory proteins expressed in transgenic swine protect swine xenografts from humoral injury. Transplant Proc 28: 758.
- 3 Dorling et al. (1996). Detection of primary direct and indirect human anti-porcine T cell responses using a porcine dendritic cell population. Eur J Immunol 26: 1378-1387.
- 4 Dorling et al. (1996) Cellular xenoresponses: Although vigorous, direct human T cell anti-pig primary xenoresponses are significantly weaker than equivalent alloresponses. Xenotransplantation 3: 149-157.
- 5 Auchincloss (1995) Why is cell-mediated xenograft rejection so strong? Xeno 3: 19.
- 6 Auchincloss (1988) Xenogeneic transplantation. Transplantation 46: 1.
- 7 Dorling et al. (1996) Cellular xenoresponses: Observation of significant primary indirect human T cell anti-pig xenoresponses using co-stimulator-deficient or SLA class II-negative porcine stimulators. Xenotransplantation 3: 112-119.
- 8 Linsley et al. (1991) CTLA-4 is a second receptor for the B-cell activation antigen B7. J Exp Med 174:561-569.
- 9 Lane et al. (1993) Expression and functional properties of mouse B7/BB1 using a fusion protein between mouse CTLA4 and human γ1. Immunology 80: 56.
- 10 Cohen (1992) Mounting a targeted strike on unwanted immune responses. Science 257:751.
- 11 Baliga et al. (1994) CTLA4Ig prolongs allograft survival while suppressing cell mediated immunity. Transplantation 58: 1082-1090.
- 12 Lenschow et al. (1992) Long term survival of xenogeneic pancreatic islet grafts induced by CTLA4-Ig. Science 257: 789-792.

- 13 Heckl-Östreicher et al. (1995) Functional activity of the membrane-associated complement inhibitor CD59 in a pig-to-human in vtiro model for hyperacture xenograft rejection. Clin. Exp. Immunol. 102:589-595.
- 14 McCurry et al. (1996) Human complement regulatory proteins expressed in transgenic swine protect swine xenografts from humoral injury. Transplant Proc 28: 758.
- 15 White et al. (1995) The control of hyperacute rejection by genetic engineering of the donor species. Eye 9: 185-189.
- 16 Yannoutsos et al. (1995) Production of pigs transgenic for human regulators of complement activation. Transplant Proc 27: 324-325.
- 17 Langford et al. (1996) Production of pigs transgenic for human regulators of complement activation using YAC technology. Transplant Proc 28: 862-863.
- 18 Bradley & Liu (1996) Target practice in transgenics. Nature Genet 14: 121-123.
- 19 Clarke (1996) The adenovirus and the egg: a new approach to transgenesis. Nature Biotech. 14:942.
- 20 Wheeler (1994) Development and validation of swine embryonic stem cells: a review. Reprod Fertil Dev 6:563-568.
- 21 Parsons et al. (1996) Cattle CTLA-4, CD28 and chicken CD28 bind CD86: MYPPPY is not conserved in cattle CD28. Immunogenetics 43: 388-391.
- 22 Dorling et al. (1996) In vitro accommodation of immortalized porcine endothelial cells. Transplantation 62:1127-1136.
- 23 Marelli-Berg et al. (1996) Major histocompatibility complex class II-expressing endothelial cells induce allospecific nonresponsiveness in naive T cells. J Exp Med 183:1603.

CLAIMS

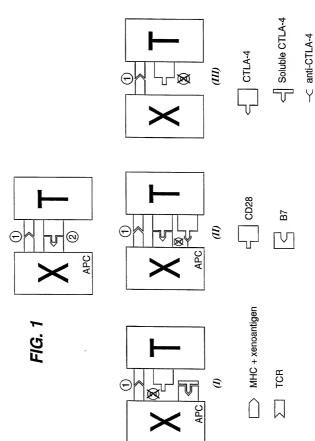
- A biological reagent capable of inhibiting T-cell mediated rejection of a xenotransplanted organ by blocking the delivery of co-stimulatory signal 2 in order to prevent the activation of xenoreactive T-cells in the recipient.
- 5 2. A method for inhibiting T-cell mediated rejection of a xenotransplanted organ, comprising blocking the delivery of co-stimulatory signal 2 in order to prevent the activation of xenoreactive T-cells in the recipient.
 - A method according to claim 2, comprising the administration to said recipient to a soluble form of the CTLA-4 protein from the xenogeneic donor organism.
- A method according to claim 3, wherein said soluble protein comprises the extracellular domain of porcine CTLA-4 fused to a human Cγ1 sequence.
 - 5. A soluble form of xenogeneic CTLA-4 for use as a medicament.
 - 6. A protein comprising the amino acid sequence SEQ ID:1
 - 7. Nucleic acid which encodes the protein according to claim 6
- 15 8. A biological reagent according to claim 1, wherein said reagent is a membrane-associated protein which can bind to CTLA-4.
 - A protein according to claim 8, comprising a single chain antibody with specificity for CTLA-4.
 - 10. Nucleic acid which encodes a protein according to claim 8 or claim 9.
- 20 11. A cell which expresses a protein according to claim 8 or claim 9 on its surface.
 - 12. Biological tissue comprising a cell according to claim 11.
 - 13. An animal comprising a cell according to claim 11 and/or biological tissue according to claim 12.
- 14. A method of transplantation comprising the step of transplanting biological tissue according to claim 12 from a donor animal into a xenogeneic recipient animal.

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- 15. A process for rendering biological tissue suitable for xenotransplantation, comprising the step of treating said biological tissue such that it expresses a protein according to claim 8 or claim 9 on the surface of its cells.
- 16. A protein according to claim 8 or claim 9, or nucleic acid according to claim 10, for use as a medicament.
- 17. The use of a protein according to claim 8 or claim 9, or of nucleic acid according to claim 10, in the preparation of a formulation for administration to a xenotransplant recipient or donor.
- 18. A biological reagent according to claim 1, wherein said reagent is a cell which expresses on its surface MHC class II of a different organism.
- 19. A cell according to claim 18, wherein said cell is a porcine cell expressing human MHC class II on its surface.
- 20. A cell according to claim 18 or claim 19, wherein said cell does not express B7 on its surface.
- 21. A cell according to claim 18 or claim 19, wherein said cell is a transfected immature dendritic cell
 - 22. Biological tissue comprising a cell according to any one of claims 18, 19, 20 or 21.
 - 23. An animal comprising biological tissue according to claim 22.
 - 24. A method of transplantation comprising the step of transplanting biological tissue according to claim 22 from a donor animal into a xenogeneic recipient animal.
 - 25. A process for rendering biological tissue suitable for xenotransplantation, comprising the step of treating said biological tissue such that it expresses xenogeneic MHC class II on the surface of its cells.
 - 26. A cell according to any one of claims 18, 19, 20 or 21, for use as a medicament.
- 25 27. The use of biological tissue according to claim 22 in the manufacture of a formulation for administering to a xenotransplant recipient.
 - 28. The use of xenogeneic MHC class II, or nucleic acid encoding xenogeneic MHC class II, in the preparation of a formulation for administering to a xenotransplant donor.

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81 KVNLTIQGLR Q•••••	141 SSGLFFYSFL	SEQ ID:1 (pCTLA4) Human CTLA4 Cattle CTLA4
71 STCTGTSTEN •I•••SG•	131 DFLLWILAAV	PIN SEQ II ••• Human ••• Cattle
61 VEDELTFLDD MGN•••••		181 CEKQFQPYFI
51 MTEVCAATYT V•••••M V••••G••M	111 VGMGNGTQIY L•I•••A••	151 161 171 181 L. TAVSLSKML KKRSPLTTGV YVKMPPTEPE CEKQFQPYFI L. *
41 VTVLRRAGSQ ••••Q•D••	101 VELLYPPPYY •••MYPPPYY	161 KKRSPLTTGV
31 GSAGKAAEVR A•P••T•• E•S••D••	91 AVDTGLYICK •M••••••	151 ITAVSLSKML L
	61 71 F VEDELTFLDD STCTGTSTEN MGN**********************************	31 41 51 61 71 81 GSAGKAAEVR VTULRRAGSQ MTEVCAATYT VEDELTFLDD STCTGTSTEN KVNLTIGGLR A•P•••T•• E•S••D•• V•••G•• 11 11 121 AVDTGLYICK VELLXZPPYY VGMGNGTQIY VIDPEPCPDS DFLLMILAAV SSGLFFYSFL •••••••••••••••••••••••••••••••••••

FIG. 3

51 TAGGACCTGG C	ATGCACGTG	171 GTGTGAGTAT ****************************	231 CGGCAGCCAG T•A•••••	291 CCTTGATGAC •••A••••T	1 GGCTGAGA •A••••G
ATGCTTGCT CTGGATTCCG GAGCCATGGG GCTTGGCTGG AGCTTACTTC TAGGACCTGG	CCCTGTACAG CTCTGTTTTC TCTTCTCTTC ATCCCTGTCT TCTCCAAAGG GATGCACGTGC.TCT.TCTCTTTCTTTCTTCTTCTTCTTCTTCTTC				301 351 TCTACATGCA CTGGCACCTC CACCGAAAAC AAAGTGAACC TCACCATCCA AGGGCTGAGA
31 GCTTGGCTGG •••CA•••A A••••	91 ATCCCTGTCT •••••••	121 151 151 161 GCCCAACCTG CAGTAGTGCT GGCCAACAGC CGGGGTGTTG CCAGCTTTGTGG TGGGG	201 221 CGAGGTCCGG GTGACAGTGC TGCGGGGGG T*****************************		331 AAAGTGAACC C
1 11 21 ATGCCTTGCT CTGGATTCCG GAGCCATGGG **********************************	B1 TCTTCTTTC G	141 GGCCAACAGC ••••G•••	201 CGAGGTCCGG T	251 TCTGTGCGG GACATATACT GTGGAGGATGGG. A.CC.CCTG AGAA	321 CACCGAAAAC ••GT•G•••T
11 CTGGATTCCG TTOA	CCCTGTACAG CTCTGTTTTCCC.TC TCT	121 GCCCAACCTG CAGTAGTGCT •••••G•••• ••••G•••• S	191 GCAAAGCTGC •••••CA•	251 TCTGTGCCGC •••••66•	311 CTGGCACCTC
ATGGCTTGCT	61 CCCTGTACAG C.TC	121 GCCCAACCTG •••••G•••• A••••G••••C	181 GGGTCTGCAG •CA•••C••	241 ATGACTGAAG G	301 TCTACATGCA ••C•TC•••

361 GCCGTGGACA •••A•••••	371 CIGGGCTCTA G.A		411 CATCTGCAAG GTGGAGCTCC TGTACCCACC ACCCTACTAT	401 TGTACCCACC	411 ACCCTACTAT G. A
421 GTGGGTATGG CCA.		431 GCAACGGGAC CCAGATTTAT GTCATTGATCAG	451 GTCATTGATC	461 CAGAACCATG	461 CAGAACCATG CCCAGATTCT
481 GATTTCCTGC ••C•••C•	491 TCTGGATCCT	501 GGCAGCAGTT T	511 AGTTCAGGGT	521 TGTTTTTTA 	531 CAGCTTCCTC TT
541 ATCACAGCTG C		TITCTTIGAG CAAAATGCTA AAGAAAAGAA GTCCTCTTAC	571 AAGAAAAGAA	571 AAGAAAGAA GTCCTCTTAC TACAGGGGTC	591 TACAGGGGTC A
601 TATGTGAAAA	611 TGCCCCGAC	621 651 AGAGCCAGAA TGTGAAAAGC AAITTCAGCC TTATTTTATT	631 TGTGAAAAGC	641 AATTTCAGCC	651 TTATTTTATT
661 CCCATCAATT GA	671 GA • •	SEQ ID: 2 (P Human CTLA4 Cattle CTLA4	SEQ ID: 2 (pcTLA4) Human CTLA4 Cattle CTLA4		

FIG. 3 (CONTD.)

331 341 351 361 SKLTVDKSRW QQGNVFSCSV MHEALHNHYT QKSLSLSPGK

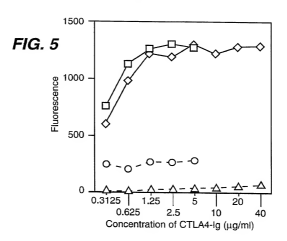
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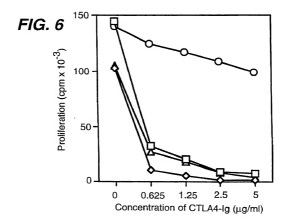
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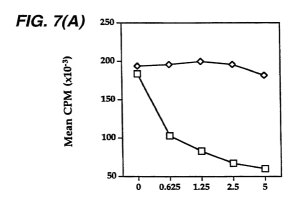
351

	-			
81	141	201	261	321
KVNLTIQGLR	KSCDKTHTCP	YVDGVEVHNA	KAKGQPREPQ	LDSDGSFFLY
71	131	191	251	271 281 291 301 321 XZTEPPSRDE LTKNQVSLTC LVKGFYPSDI AVEWESNGQP ENNYKTTPPV LDSDGSFFLY
STCTGTSTEN	D <u>GGSGGAA</u> EP	HEDPEVKFNW	LPAPIEKTIS	
61	121	181	241	301
VEDELTFLDD	VIDPEPCPDS	EVTCVVVDVS	EYKCKVSNKA	AVEWESNGQP
51	111	171	231	291
MTEVCAATYT	VGMGNGTQIY	KDTLMISRTP	VLHQDWLNGK	LVKGFYPSDI
	101	161	221	281
VTVLRRAGSQ	VELLYPPPYY	PSVFLFPPKP	STYRVVSVLT	LTKNQVSLTC
31	91	151	211	271
GSAGKAAEVR	AVDTGLYICK	PCPAPELLGG	KTKPREEQYN	VYTLPPSRDE
	41 51 61 71 VIVERRAGSQ MTEVCAATYT VEDELTFLDD STCTGTSTEN KVNLTIQ	31 41 81 GSAGKAAEVR VTVLRRAGSQ MTEVCAATYT VEDELTFLDD STCTGTSTEN KVNLTIQGLR 91 101 121 131 AVDTGLYICK VELLYPPPYY VGMGNGTQIY VIDPEPCPDS DGGSGGAAEP KSCDKTHTCP	31 41 51 61 71 81 GSAGKAAEVR VTVLRRAGSQ MTEVCAATYT VEDELTFLDD STCTGTSTEN KVNLTIQGLR 91 101 111 121 131 141 AVDTGLYICK VELLYPPPYY VGMGNGTQIY VIDPEPCPDS DGGSGGAAEP KSCDKTHTCP 151 161 171 181 201 PCPAPELLGG PSVFLFPPRP KDTLMISRTP EVTCVVUDVS HEDPEVKFNM YVDGVEVHNA	31 41 81 GSAGKAAEVR VTVLRRAGSQ MTEVCAATYT VEDELTFLDD STCTGTSTEN KVNLTIQGLR 91 101 111 121 141 AVDTGLYICK VELLYPPPYY VGMGNGTQIY VIDPEPCPDS DGGSGGAAEP KSCDKTHTCP 151 161 171 181 201 PCPAPELLGG PSVFLFPPKP KDTLMISRTP EVTCVVVDVS HEDPEVKFNW YVDGVEVHNA 211 221 231 231 241 251 261 KTKPREEQYN STYRVVSVLT VLHQDWLNGK EYKCKVSNKA LPAPIEKTIS KAKGQPREPQ

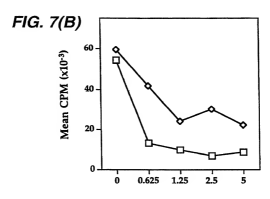
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Concentration of CTLA4-Ig (µg/ml)



Concentration of CTLA4-Ig (µg/ml)

721 GT

FIG. 8

. 4	. (1)		. 64			. (1)	. ()	. (1)	
rcigg	ragtg	CGCTG	SACTA	ACTTC	SCTCC	AATCA(CCGGT	rccta(
71	151. CTAT	231.	311.	391. GTGC	471.	551. GAAT	631	711. ACCG1	
TGTGC	CTCAG	ccaag	ATTT	CGGTA	CIGGA	TATAG	CAGTG	AGCTG	
61 TCTCC	141 TGGGT	221 CAATT	301 GTCGT	381 TCTGG	461 TTGTT	541 TCATC	621 GCCAT	701 GACCA	
SAGAC	rggAG	GAGA	GCTG	TGGC	TCTC	ACTCC	CCTG	GAGG	
51	131 GGGGC	211 TCTCC	291 GCAAG	371	451	531 cccaa	611	691 TCGGC	
99999	SGGAA	ACCA	CTGT	CAGG	SAGGG	22552	ACCTC	SGTAT	
41	121 CTCCAG	201	281 GTATT	361 GCGGT7	441	521 AGGAAG	601 CTGGC	681	
STACA	cage	4666C	SCCGT	regag	20000	TCCC	AAGT	TGAC	
31 GCTTG	111 GTCCG	191 CGTGA	271 ACACG	351	431 TGGGA	511	591 GGCTC	671 ATGGG	
GGAG	SCTGG	SACTC	GAGG	CTCG	CGTC	TACC	CTCT	CAGC	
21 TCTGGG	101 CATGAC	181	261 AGAGCO	341	421	501	581	661	
regae	TATGC	TACT	SCCTG	TGGT	sccac	TGTA	CTGA	TATT	
11 GCTGG	91	171	251 GAACAG	331 GTACC	411	491 TAATT	571 GGGTC	651 GCTGA1	
STGCA	81 91 101 111 121 131 141 151 17 praccrtta gcagcigeag cargagcige ciccage ciccageaa gggcigeag iggercicae cartaafige		CAAAT	CCAAG	TGCTG	GAAG	TCAG	641 651 661 671 681 691 711 711 711 711	
1	81	161 171 181 191 201 211 221 231 ragiggiggt agcacatact acgasactc cgicaaaggc cgettcacca ictccagaga caattccaag aacacgeter	241 251 261 271 291 311 311 311 311 311 311	321 331 341 351 361 371 381 391 rggggccaag gracctag gractag gra	401 411 421 431 441 451 461 471	481 491 501 511 521 531 541 551	561 571 581 591 601 611 621 631 cgeccercas gegrecers gecareast gegrecest	641	

FIG. 9

81....... 91....... 101...... 111...... 121....... 131....... 141...... 151....... 151....... 150...... 150.....

161...... 171...... 181...... 191...... 201...... 211...... 221...... 231...... IGSNIYYYWYQ QLPGTAFKLL IYRNNQRPSG VPDRFSGSKS GTSASLAISG IRSEDEADYY CAAWDDSLVF GGGTKLITVLG

3 3-2-7 Ġ

Stimulators (x10 -4)

c.p.m. (x 10-3)

FIG. 13

FIG. 10

	1	0 	20	30 	40 	50 	60
M1 sFv M3 sFv M19 sFv M24 sFv	CATGG-CCC CATGG-CCC	AGGTGCAC AGGTGCAC	CTGCAGGAG CTGGTGCAG	TCGGGGCCCAG TCTGGGGGCTG	GACTGGTGA AGGTGAAGA	AGCCTGGGG AGCCTTGGGA GGCCTGGGGG AGCCCGGGGA	GACCCTGTC CTCAGTGAA
	70 	80 	90 	100) 11	0 12 	0
M1 sFv M3 sFv M19 sFv M24 sFv	CCTCACCTC	CACTGTC	CTGGTGGCT CTGGATACA	CCGTCAGCAG	TGGTAGTTA CTA	CCATGAGO CTACTGGAGO CTATATGCAO CTGGATCGGO	TGGATCCGG TGGGTGCGA
	130	140 	150	160 	170 	180 	190
M1 sFv M3 sFv M19 sFv M24 sFv	CAGCCCCCA	GGGAAGG(GGACAAG	SACTOGAGTG SGCTTGAGTG	GATTGGGT GATGGGAATZ	PAT-ATCTAT	AGTGGTGGTA TACAGTGGGA AGTGGTGGTA GGTGACTCTG	GCACCAACT GCACAAGCT
	200 	2:	LO 2 	20 2	230	240	250
M1 sFV M3 sFV M19 sFV M24 sFV	ACGCAGACT ACAACCCCT ACGCACAGA	CCCTCAA CCCTCAA	GGCCGGTTC GAGTCGAGTC GGGCAGAGTC	ACCATCTCC: ACCATATCAC ACCATGACC:	AGAGACAATT STAGACACGT AGGGACACGT	240 CCAAGAACAC CCACGAGCAC CCACGAGCAC	GCTGTATCT GTTCTCCCT 'AGTCTACAT
M3 sFv M19 sFv	ACGCAGACT ACAACCCCT ACGCACAGA	CCCTCAA CCCTCAA	GGCCGGTTC GAGTCGAGTC GGGCAGAGTC	ACCATCTCC: ACCATATCAC ACCATGACC:	AGAGACAATT STAGACACGT AGGGACACGT	CCAAGAACAC CCAAGAACCA CCACGAGCAC	GCTGTATCT GTTCTCCCT 'AGTCTACAT
M3 sFv M19 sFv	ACGCAGACI ACAACCCCI ACGCACAGE ACAGCCCGI 260 GCAAATGAI GAAGCTGAC GGAGCTGAC	COGTIGAA COCTICAA AGITTOCA 270 270 ACAGCCTIG	GGGCCGGTTC GAGTCGAGTC GGGCAGAGTC 280 AGAGCCGAGGAGAGCCAGGAGAGAGAGCCGAGGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAG	ACCATCTCC: ACCATCTCC: ACCATGACC: ACCATCTCAC 290 ACCACGGCCC: ACCACGGCCC: ACCACGGCCC: ACCACGGCCC: ACCACGGCCCC: ACCACGGCCCC: ACCACGGCCCC: ACCACGGCCCC:	AGAGACAATTT STAGACACGT AGGGACACGT GCCGACAAGT 300 INSTATTACTG INSTATTACTG	CCAAGAACAC CCAAGAACCA CCACGAGCAC CCATCAGCAC	CONGRATOR GETTACAT AGRICIACAT COGCOTACCT 320
M3 SFV M19 SFV M24 SFV M1 SFV M3 SFV M19 SFV	ACGCAGACTI ACAACCCCTI ACGCACAGI ACAGCCCGTI 260 	COGTIGAA COCTICAA AGITTOCA 270 270 ACAGCCTIG	GGGCCGGTTC GAGTCGAGTC GGGCAGAGTC 280 AGAGCCGAGGAGAGCCAGGAGAGAGAGCCGAGGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAG	ACCATCTCC: ACCATCTCC: ACCATGACC: ACCATCTCAC 290 ACCACGGCCC: ACCACGGCCC: ACCACGGCCC: ACCACGGCCC: ACCACGGCCCC: ACCACGGCCCC: ACCACGGCCCC: ACCACGGCCCC:	AGAGACAATTT STAGACACGT AGGGACACGT GCCGACAAGT 300 INSTATTACTG INSTATTACTG	CCAAGAACAC CCAAGAACAC CCACGAGCAC CCATCAGCAC 310 TGCAAGAGCT TGCAAGAATG	CONGRATOR GETTACAT AGRICIACAT COGCOTACCT 320

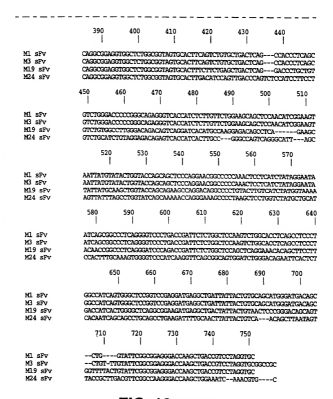
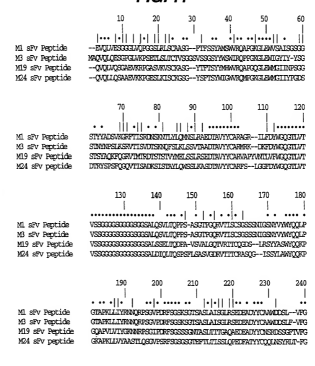


FIG. 10 (CONTD.)

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FIG. 11



M1 sFv Peptide M3 sFv Peptide M19 sFv Peptide M24 sFv peptide GGIKLIVLG
GGIKLIVLGAA
GGIKLIVLGAA
GGIKLIVLG

FIG. 12

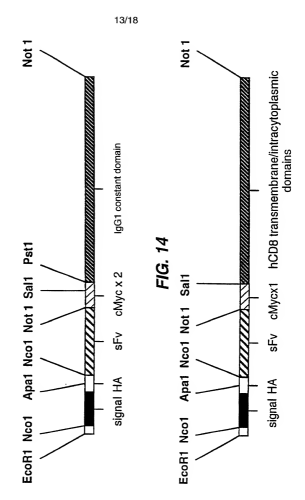


FIG. 15(A)

-65 AGCTTCAGGA TCCTGAAAGG TTTTGCTCTA CTTCCTGAAG ACCTGAACAC
-15 CGCTCCCATA AAGCCATGGC TTGCCTTGGA TTTCAGCGGC ACAAGGCTCA
36 GCTGAACCTG GCTACCAGGA CCTGGCCCTG CACTCTCCTG TTTTTTCTTC
86 TCTTCATCC TGTCTTCTGC AAAGCAATGC ACGTGGCCCA GCCTGCTGTG
136 GTACTGGCCA GCCCCAGGG CATCGCCAGC TTTGTGTTGT AGTAGCATC
ACGCCAAA GCCACTGAGG TCCGGGTGAC AGTGCTCTGG CAGGCTGACA
236 GCCAGGTGAC TGAAGTCTGT GCGGCAACCT ACATGATGGG GAAATGAGTTG
286 ACCTTCCTAG ATGATTCCAT CTGCACGGG ACCTCCAGTG GAAATCAAGT
336 GAACCTCACT ATCCAAGGAC TGAGGGCCAT ACTACCTGGG CATAGGCAAC
436 GCAAGGTGGA GCTCATGTAC CCACCGCCAT ACTACCTGGG CATAGGCAAC
436 GCAACCTCAC TTTATGTAAT TGATCCAGAA CCGTGCCCAG ATTCTGACTT
486 CCTCCTCTGA ATCCTTGCAG CAGTTAGTTC GGGGTTGTTT TTTTATACCT
536 TTCTCCCCAC AGCTGTTCT TTGAGCAAAA TGCTAAAGAA AAGAAGCCCT
536 AAAGCAATT CAGCCTTATT TTATTCCCAT CAAATGAGGAA TT

FIG. 15(B)

-30 -20 -10 1 11 21 MACLGFQRHK AQLNLAIRTW PCTLLFFLLF 1PVFCKAMHV AQPAVVLASS RGIASFVCEY

31 41 51 61 71 81 ASPGKATEVR VTVLRQADSQ VTEVCAATYM MGNELTFLDD SICTGTSSGN QVNLTIQGLR

91 101 111 121 131 141 AMDIGLYICK VELMYPPPYY LGIGNG $\underline{T}Q$ IY VIDPEPCPDS DFLLWILAAV SSGLFFYSFL

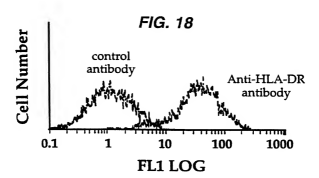
151 161 171 181 LTAVSLSKML KKRSPLTTGV YVKMPPTEPE CEKQFQPYFI PIN

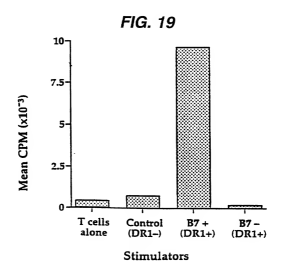
FIG. 16

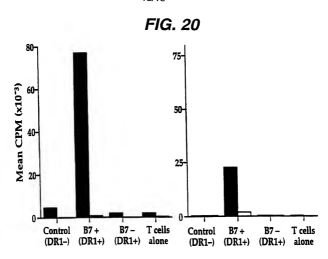
-36	AAGCTTCGAG	CCAAGCAGCG	TCCTGGGGAG	CGCGTC <u>ATG</u> G	CCTTACCAGT
15	GACCGCCTTG	CTCCTGCCGC	TGGCCTTGCT	GCTCCACGCC	GCCAGGCCGA
65	GCCAGTTCCG	GGTGTCGCCG	CTGGATCGGA	CCTGGAACCT	GGGCGAGACA
115	GTGGAGCTGA	AGTGCCAGGT	GCTGCTGTCC	AACCCGACGT	CGGGCTGCTC
165	GTGGCTCTTC	CAGCCGCGCG	GCGCCGCCGC	CAGTCCCACC	TTCCTCCTAT
215	ACCTCTCCCA	AAACAATCCC	AAGGCGGCCA	AGGGGCTGGA	CACCCAGCGG
265	TTCTCGGGCA	AGAGGTTGGG	GGACACCTTC	GTCCTCACCC	TGAGCGACTT
315	CCGCCGAGAG	AACGAGGGCT	ACTATTTCTG	CTCGGCCCTG	AGCAACTCCA
365	TCATGTACTT	CAGCCACTTC	GTGCCGGTCT	TCCTGCCAGC	GAAGCCCACC
415	ACGACGCCAG	CGCCGCGACC	ACTAACACCG	GCGCCCACCA	TCGCGTCGCA
465	GCCCCTGTCC	CTGCGCCCAG	AGGCGTGCCG	GCCAGCGGCG	GGGGGCGCAG
515	TGCACACGAG	GGGGCTGGAC	TTCGCCTGTG	ATATCTACAT	CTGGGCGCCC
565	CTGGCCGGGA	CTTGTGGGGI	CCTTCTCCTG	TCACTGGTTA	TCACCCTTTA
615	CTGCAACCAC	'AGGAACCGAA	GACGTGTTTG	CAAATGTCCC	CGGCCTGTGG
665	TCAAATCGGG	AGACAAGCCC	AGCCTTTCGG	CGAGATACGT	CTAACCCTGT
715	GCAACAGCC	CTACATGAAT	TCC		

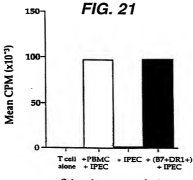
FIG. 17 **B7+ IPEC** hCTLA4-Ig Mean Relative fluorescence Intensity mCTLA4-Ig 2 0.001 0.01 0.1 1 10 100 **B7 - IPEC** 6 hCTLA4-Ig 2 mCTLA4-Ig 0.001 0.01 0.1 1 10 100

CTLA4-Ig concentration (µg/ml)









Stimulator population

COMBINED DECLARATION FOR PATENT APPLICATION AND POWER OF ATTORNEY T International Applications (Includes Reference to Pi

ATTORNEY DOCKET NUMBER 2292/0H795

As a below named inventor, I hereby declare that:

My residence, post office address and citizenship are as stated below next to my name.

I believe I am the original, first and sole inventor (if only one name is listed below) or an original, first and joint inventor (if plural names are listed below) of the subject matter which is claimed for and which a patent is sought on the invention entitled:

IMMUNOSUPPRESSION BY BLOCKING T CELL CO-STIMULATION SIGNAL 2(B7/CD28 INTERACTION)

the spec	cification of which (check only one item belo
[X]	is attached hereto.
O	was filed as United States application
	Serial No.
	on
	and was amended
	on (if applicable).

ESS June was filed as PCT international application (X3

Number PCT/GB99/01350 on 30 April 1999 and was amended under PCT Article 19 __ (if applicable)

I hereby state that I have reviewed and understand the contents of the above-identified specification, including the claims as amended by any amendment referred to above.

I acknowledge the duty to disclose information which is material to the examination of this application in accordance with Title 37, Code of Federal Regulations, §1.56(a).

I hereby claim foreign priority benefits under Title 35, United States Code, §119 of any foreign application(s) for patent or inventor's certificate or of any PCT international application(s) designating at least one country other than the United States of America listed below and have also identified below any foreign application(s) for patent or inventor's certificate or any PCT international application(s) designating at least one country other than the United States of America filed by me on the same subject matter having a filing date before that of the application(s) of which priority is claimed:

PRIOR FOREIGN/PCT APPLICATION(S) AND ANY PRIORITY CLAIMS UNDER 35 U.S.C. 119:

COUNTRY (d PCT minuse PCT)	APPLICATION NUMBER	DATE OF FILING (day, month, year)	, PRIORIT UNDER	Y CLAIMED If U.S.C. 119
Great Britain	9809280,2	30 April 1998	[X] YES	[]wo
			[] YES	Ом []
	·		Q YES	[] NO
			[] YES	[] NO
			[] YES	() NO

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(Japuary 1991)

JŠ

Combined Declaration for Patent Application and Power of Attorney (Continued)

(Includes Reference to PCT International Applications)

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I hereby claim the Pecefit under Title 35. United States Code, \$120 of any United States applications(s) expension of the States and Pecefit and States of America that is large its often below and, insoft as the subject matter of each of the claims of this applications is not disclosed in hardfoor prior application(s) in the manner provided by the first paragraph of Title 35, United States Code \$11.2. I acknowledge the duty to disclose material information as defined in Title 37, Code of Pederal Regulations, \$1.56(a) which occurred between the filing date of the prior application(s) and the national or PCT international filing date of this applications.

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35 U.S.C. 120:

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POWER OF ATTORNEY: As a named inventor, I hereby appoint the following attorney(s) and/or agents to prosecute this application and transact all business in the Patent and Trademark Office connected therewith. Morris Relon #15,108, Gordon D. 169; #19,168, William F. Dudine, Jr. #20,569, Michael J. Sweedler #19,937, S. Peter Ludwig #25,351, Paul Fields #20,298, Joseph B. Lerch #26,353, Melvin C. Garner #26,227.2. Ethan Horwitz #27,646, Severty B. Goodwin #26,417, Adda C. Goognis #29,414, Martin E. Goldfrid #20,889, Bert J. Lewen 19,407, Henry Stemberg #22,408, Peter C. Schechter #31,662, Robert Schaffer #31,194,Robert C. Sulfiken, Jr. #20,499, and Joseph R. Robinson #33,448

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100	POST OFFICE ADDRESS	POST OFFICE ADDRESS Hammersmith Hospital	CITY London W12 ONN	STATE & 2F CODE/COUNTRY United Kingdom
2.4	FULL NAME DF INVENTOR	DORLING 2 -00	FIRST GIVEN NAME Anthony	SECOND GIVEN NAME
0	RESIDENCE & CITIZENSHIP	London W12 ONN GBX	STATE OR FOREIGN COUNTRY United Kingdom	COUNTRY OF CITIZENSHIP United Kingdom
2	POST OFFICE ADDRESS	POST OFFICE ADDRESS Hammersmith Hospital	CITY London W12 ONN	STATE & ZIP CODE/COUNTRY United Kingdom
2	OF INVENTOR	FAMILY NAME	FIRST GIVEN NAME	SECOND GIVEN NAME
0	RESIDENCE & CITIZENSHIP	CITY	STATE OR FOREIGN COUNTRY .	COUNTRY OF CITIZENSHIP
3	POST OFFICE ADDRESS	POST OFFICE ADORESS	CITY	STATE & 2IP CODE/COUNTRY

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	lew	A-Doney	
DAYE 13- 1	00	DATE 13/4/00	DATE